



Digestion pattern of reovirus outer capsid protein $\sigma 3$ determined by mass spectrometry

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Abstract

Reovirus is an enteric virus comprising eight structural proteins that form a double-layered capsid. During reovirus entry into cells, the outermost capsid layer (composed of proteins $\sigma 3$ and $\mu 1C$) is proteolytically processed to generate an infectious subviral particle (ISVP) that is subsequently uncoated to produce the transcriptionally active core particle. Kinetic studies suggest that protein $\sigma 3$ is rapidly removed from virus particles and then protein $\mu 1C$ is cleaved. Initial cleavage of $\mu 1C$ has been well described and generates an amino (N)-terminal δ peptide and a carboxyl (C)-terminal ϕ peptide. However, cleavage and removal of $\sigma 3$ is an extremely rapid event that has not been well defined. We have treated purified reovirus serotype 1 Lang virions with a variety of endoproteases. Time-course digestions with chymotrypsin, Glu-C, pepsin, and trypsin resulted in the initial generation of two peptides that were resolved in SDS–PAGE and analyzed by in-gel tryptic digestion and MALDI-Qq-TOFMS. Most tested proteases cut $\sigma 3$ within a “hypersensitive” region between amino acids 217 and 238. In addition, to gain a better understanding of the sequence of subsequent proteolytic events that result in generation of reovirus subviral particles, time-course digestions of purified particles were performed under physiologic salt conditions and released peptide fragments ranging from 500 to 5000 Da were directly analyzed by MALDI-Qq-TOFMS. Trypsin digestion initially released a peptide that corresponded to the C-terminus of $\mu 1C$, followed by a peptide that corresponded to amino acids 214–236 of the $\sigma 3$ protein. Other regions of $\mu 1C$ were not observed until protein $\sigma 3$ was completely digested. Similar experiments with Glu-C indicated the hypersensitive region of $\sigma 3$ was cut first when virions were treated at pH values of 4.5 or 7.4, but treatment of virions with pepsin at pH 3.0 released different $\sigma 3$ peptides, suggesting acid-induced conformational changes in this outer capsid protein. These studies also revealed that the N-terminus of $\sigma 3$ is acetylated.

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Introduction

A large number of viruses (e.g., caliciviruses, coronaviruses, many enteric picornaviruses, and reoviruses) are primarily transmitted from host to host by the fecal–oral route. Thus, the predominant route of host infection by such viruses is enteric. During a gastrointestinal infection, such viruses must go through the stomach and endure an acidic pH that can be as low as 0.82 (Vander et al., 1994). Successful replication of viruses, similar to other obligate intracellular pathogens, requires that the pathogenic organism

gain entry into susceptible host cells. Initial steps in such a process involve recognition of the host cell by appropriate viral proteins. After attachment of the virus to a suitable receptor, the virus must be internalized. While some details of how viruses are internalized remain controversial, there is mounting evidence that entry of many viruses is mediated by conformational changes that take place in virus structural proteins. Some of these changes appear to involve exposure of the virus to acidic pH and others appear to involve proteolysis of external virion proteins (White and Fenner, 1994; Young, 2001).

The mammalian orthoreoviruses (reovirus) are the prototypic members of the nonfusogenic *Reoviridae*, enteric viruses that consist of a genome of double-stranded (ds) RNA encased in multiple concentric protein capsids. Reo-

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viruses are constructed from eight structural proteins arranged as two concentric capsids; an outer capsid that surrounds an inner capsid “core” (reviewed in Nibert and Schiff, 2001). The outer capsid is composed of up to 36 copies of the cell attachment protein $\sigma 1$ (Larson et al., 1994), 600 copies of protein $\sigma 3$ (Dryden et al., 1993), and 600 copies of protein $\mu 1$, which occurs in the viral capsid both as $\mu 1$ and its cleavage product $\mu 1C$ (Nibert and Fields, 1992). The inner core, which is the transcriptionally active structure that produces progeny capped and methylated mRNA from the genomic dsRNA (Yeager et al., 1996), comprises two major proteins (120 copies of $\lambda 1$ and 150 copies of $\sigma 2$; Reinisch et al., 2000), and two minor proteins (12 copies of $\lambda 3$ and about 20 copies of $\mu 2$; Coombs, 1998). Sixty copies of an additional protein ($\lambda 2$) form pentameric spike structures at each of the virion’s icosahedral vertices (Ralph et al., 1980; Metcalf et al., 1991; Dryden et al., 1993) that bridge the inner and outer capsid layers.

Entry of reovirus into cells is accompanied by significant alterations in the virus outer capsid, which results in the generation of an infectious (or intermediate) subviral particle (ISVP). The virus attaches to the receptor using its cell attachment protein $\sigma 1$ (Lee et al., 1981). Binding of the virion to the receptor leads to a conformational change in the viral capsid as demonstrated by reversible alterations in pepsin sensitivity (Fernandes et al., 1994). Such an altered conformational state may be necessary for viral entry and subsequent uncoating.

Once attached, the virion appears to be taken up into endosomes (Borsa et al., 1979; Georgi et al., 1990; Rubin et al., 1992). An influx of H^+ ions, mediated by proton pumps present on the endosomal membrane, creates an acidic environment, which, in conjunction with the presence of acid-dependent proteases such as cathepsins (Sherry et al., 1996; Baer and Dermody, 1997), is believed to proteolytically process the virion into subviral particles such as ISVPs. Upon exposure of the ISVP to the cytosol it is subsequently converted into the core particle. Evidence also suggests that ISVPs may be formed extracellularly by proteolysis of the virion with proteases such as pepsin in the stomach, or chymotrypsin or trypsin in the intestinal lumen (Bodkin et al., 1989; Vander et al., 1994), and that these ISVPs can directly cross through the membrane, bypassing endocytic uptake (Borsa et al., 1979; Sturzenbecker et al., 1987).

Proteolytic processing of the outer capsid also is achieved *in vitro* by treatment of virions with any of a variety of proteases, including the alkaline proteases trypsin and chymotrypsin (Joklik, 1972) and acidic proteases such as cathepsins (Ebert et al., 2002; Jane-Valbuena et al., 2002). Such *in vitro* proteolysis, mediated either by trypsin or by chymotrypsin, is very rapid. Most $\sigma 3$ is removed from particles within 1 min of protease treatment (Nibert et al., 1995). Thus, details of $\sigma 3$ cleavage have not been completely delineated, although a region near amino acid residues 210–242 was very recently shown to initially be cleaved by various alkaline and acidic proteases (Ebert et

al., 2002; Jane-Valbuena et al., 2002). Subsequent cleavage of $\sigma 3$, which appears to be a prerequisite for removal of this protein from the particle and conversion of the virion into the ISVP (Jane-Valbuena et al., 2002), has not yet been defined.

To delineate early steps in virus entry mediated by outer capsid proteolysis, we have determined the kinetics of $\sigma 3$ digestion in purified intact reovirus type 1 Lang virions. Virions were digested with various proteinases for various periods of time. Products then were analyzed by matrix-assisted laser desorption ionization quadrupole/time-of-flight mass spectrometry (MALDI-Qq-TOFMS). These studies indicate the $\sigma 3$ hypersensitive region is initially cut by a large number of endoproteases under a wide range of pH conditions (from acidic to neutral), the nature of the blocking group at the N-terminus of $\sigma 3$, and the sequence and kinetics of $\sigma 3$ proteolysis during subsequent virion uncoating events.

Results

Proteolysis of reovirus virions alters outer capsid proteins $\sigma 3$ and $\mu 1$ to generate subviral particles

Treatment of purified reovirus virions with 200 $\mu g/ml$ α -chymotrypsin, conditions generally used for *in vitro* generation of subviral particles such as the ISVP and core (Joklik, 1972; Coombs, 1998), resulted in the rapid removal of $\sigma 3$ protein from virions (Fig. 1A). Other reports indicate similar results are obtained when virions are treated with similar concentrations of trypsin (Nibert et al., 1995). In each of these studies, all detectable $\sigma 3$ was digested within 1–2 min. Under these conditions, digestion of $\mu 1C$ into δ and ϕ occurs shortly after $\sigma 3$ has been removed, implying that $\sigma 3$ occupies an outermost layer within the virus particle that covers accessible sites in $\mu 1C$, consistent with structural interpretations from electron cryomicroscopic (Dryden et al., 1993) and X-ray crystallographic (Liemann et al., 2002) image reconstructions.

Reovirus outer capsid virion protein $\sigma 3$ is initially cut by a variety of proteases in one region

To slow the digestion of $\sigma 3$ and attempt to identify regions of outer capsid protein $\sigma 3$ that are initially cut, and presumably exposed on the surface of the intact virion, we then digested highly purified virions with a significantly lower amount of enzyme for shorter and comparable amounts of time and analyzed resulting products. Treatment of virions with 10 $\mu g/ml$ α -chymotrypsin at 37°C for as little as 6 s resulted in the appearance of two smaller polypeptides with apparent molecular weights of 27 and 12.5 kDa, as resolved in SDS-PAGE (Fig. 1B, arrowheads). The sum of these two polypeptide fragment molecular weights is close to the molecular weight of the 41 kDa $\sigma 3$

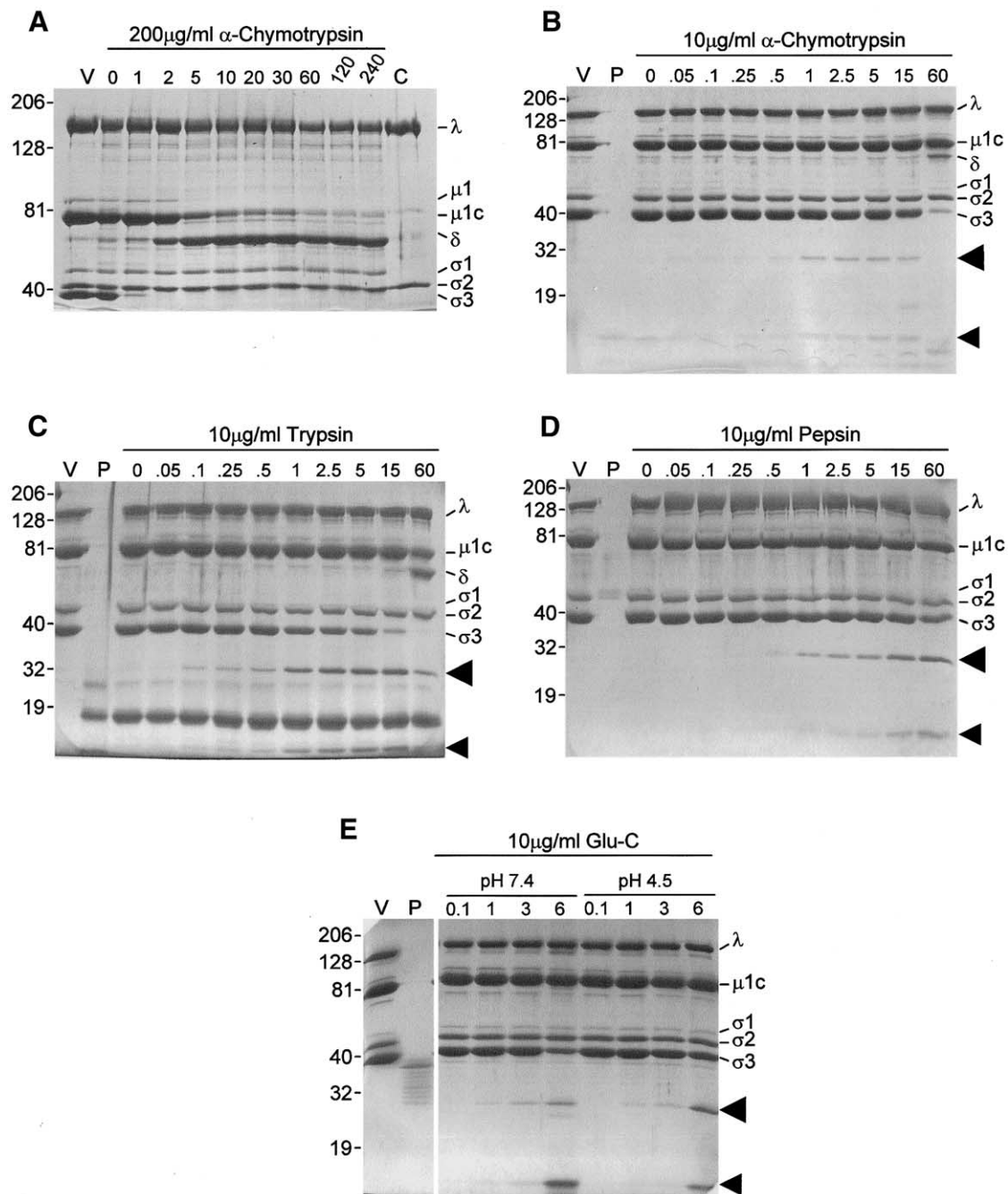


Fig. 1. Kinetics of virus digestion. Aliquots of gradient-purified T1L virions were diluted to concentrations of $7.3\text{--}8.6 \times 10^{11}$ particles/ μL in appropriate buffers (pH 7.4 for α -chymotrypsin, trypsin, and *S. aureus* V8 (Glu-C); or pH 4.5 for Glu-C; or pH 3.0 for pepsin), digested with either (A) 200 $\mu\text{g}/\text{ml}$ α -chymotrypsin, or (B) 10 $\mu\text{g}/\text{ml}$ α -chymotrypsin, (C) 10 $\mu\text{g}/\text{ml}$ trypsin, (D) 10 $\mu\text{g}/\text{ml}$ pepsin, or (E) 10 $\mu\text{g}/\text{ml}$ Glu-C for the indicated periods of time (minutes for (A–D); hours for (E)) at 37°C. Reactions were stopped by chilling and addition of phenylmethylsulfonyl fluoride to a final concentration of 5 mM to α -chymotrypsin reactions, by addition of soybean trypsin inhibitor to a final concentration of 125 $\mu\text{g}/\text{ml}$ to trypsin reactions, by adjustment of pepsin reactions to a pH of 7.5, or by chilling Glu-C reactions. One-quarter volume of 5 \times electrophoresis sample buffer was added and peptides from about 1×10^{11} particles were resolved in either a 4–16% exponential gradient (A) or a 6–16% linear gradient (B–E) mini-SDS-PAGE ($8 \times 10 \times 0.04$ cm) at 150–180 V for 60 min. Gels were fixed and stained with Coomassie brilliant blue R-250. V: gradient-purified virions; P: 150 ng of indicated protease; C: gradient-purified core particles, separately prepared. Molecular weights, determined from coelectrophoresis of Kaleidoscope Rainbow Marker, are indicated to left of each gel. Virion proteins (λ , μ , and $\sigma 1\text{--}\sigma 3$), $\mu 1\text{C}$ digestion product δ , and potential $\sigma 3$ digestion products (indicated by arrowheads) are indicated to the right of each gel.

protein, suggesting the two peptides result from a single cleavage of the protein. Longer digestions lead to the appearance of additional smaller peptides and the eventual generation of a peptide profile similar to that of ISVPs (Fig.

1B, rightmost lane). Treatment of purified virions with low concentrations of other proteases (trypsin, Glu-C, and pepsin) at various pH values for suitable periods of time also led to the generation of two polypeptides (Fig. 1C–E, ar-

rowheads), whose aggregate molecular weights added to about 41 kDa. To identify each of the polypeptides generated by each of the proteases and determine precisely where $\sigma 3$ was cut by each of them, we resolved larger aliquots of briefly digested virions in preparative gels, excised the 27- and 12.5-kDa bands, and subjected each to in-gel tryptic digestion. Digests then were extracted from the gel matrix as described under materials and methods and examined by MALDI-Qq-TOFMS (i.e., peptide mass mapping) (Fig. 2B–E). All peaks detected were checked against a database of predicted $\sigma 3$ and trypsin peptide masses to assign peaks to appropriate $\sigma 3$ peptides (Table 1) and the amino acid sequences of some peaks were examined by tandem mass spectrometry to corroborate amino acid assignments. For each of the larger 27-kDa polypeptides, we were able to identify a number of $\sigma 3$ peptides, including those corresponding to amino acids 23–34, 123–143, and 203–208 (Fig. 2F, bars above sequence scale line). Each of the smaller 12.5-kDa polypeptides contained several $\sigma 3$ peptides, including those corresponding to amino acids 248–259, 297–310, and 313–326 (Fig. 2F, bars below sequence line). In addition, depending upon the initial protease used prior to gel purification, additional tryptic fragments were identified that allowed us to determine the exact cleavage sites used by each protease. For example, the large 27-kDa polypeptide generated by chymotrypsin digestion contained a tryptic fragment with an m/z value of 2918.359, predicted to correspond to amino acids 214–238 and the small 12.5-kDa polypeptide contained a tryptic fragment with an m/z value of 1069.657, predicted to correspond to amino acids 239–247. In addition, the large 27-kDa polypeptide contained no tryptic fragments that would correspond to amino acids from residue 239 to the C-terminus and the small 12.5-kDa polypeptide contained no tryptic fragments that would correspond to amino acids from the N-terminus to residue 238. Analyses of these fragments accounted for 220 of the 365 predicted amino acids in $\sigma 3$ (60.3% coverage). We initially did not observe fragments that would correspond to either the N-terminus (residues 1–22) or the C-terminus (348–365). However, these terminal fragments were detected in direct MALDI-Qq-TOFMS measurements of longer digestions (below), suggesting they were present within the 27- and 12.5-kDa polypeptides but not effectively eluted from the gel matrix. Taken together, these observations indicate that the initial cut made in $\sigma 3$ by α -chymotrypsin occurs between Tyr₂₃₈ and Arg₂₃₉ (Fig. 2G, H), also as very recently determined (Jane-Valbuena et al., 2002). The identities of many of these tryptic fragments were corroborated by tandem mass spectrometry (MS) (see below). Similar analyses of the large and small peptides generated by other proteases indicated that trypsin cut predominantly between Arg₂₃₆ and Ala₂₃₇, although, in a small proportion of $\sigma 3$ molecules, the initial trypsin cut was between Arg₂₅₉ and Ala₂₆₀ (which is consistent with the appearance of a doublet in the gel (Fig. 2A)), and Glu-C cut predominantly between Glu₂₁₇ and Trp₂₁₈, irrespective of

whether the reaction was performed at pH values of 7.4 or pH 4.5, which is consistent with the slightly faster migration of the Glu-C-generated large peptides (Fig. 2A). There were some differences in the distribution of tryptic peptides identified after Glu-C digestion under different pH conditions, suggesting some minor acid-induced conformational changes in $\sigma 3$. No peaks corresponding to amino acids C-terminal of residue 214 were found in the 27-kDa polypeptide generated by Glu-C digestion at pH 7.4 (Table 1). However, small peaks corresponding to tryptic fragments 214–236 (m/z 2647.335) and 209–236 (m/z 3290.522) were observed in spectra generated from 27-kDa polypeptides purified after Glu-C digestion at pH 4.5 (Table 1), suggesting that, in some particles treated at this acidic pH, Glu-C cut between Glu₂₄₁ and Leu₂₄₂. Analyses of tryptic fragments in the large and small pepsin-generated peptides indicated that this protease initially cut $\sigma 3$ at one of two possible sites, and with apparently nearly equal efficiency. Thus, some $\sigma 3$ molecules were initially cut between Ala₂₃₇ and Tyr₂₃₈, whereas other $\sigma 3$ molecules were cut between Thr₂₄₄ and Pro₂₄₅.

Kinetic digestion analyses of reovirus outer capsid proteins $\sigma 3$ and $\mu 1c$

Major advantages of the MALDI-Qq-TOFMS include the capacity to rapidly analyze complex mixtures of peptides in physiologic salt solutions without the need for additional purification methodologies which might lead to sample loss. In addition, whereas gel analyses required visualization of nanogram quantities of peptide, the approximately 100-fold greater sensitivity of the MALDI-Qq-TOFMS instrument might reveal early rare events undetectable by SDS-PAGE strategies. Thus, to perform more complete analyses of the cascade of proteolytic events that take place in multiple viral capsid proteins during digestion with various proteases at different pH values, we treated purified reovirus particles with proteases used above and analyzed released peptide fragments directly by MALDI-Qq-TOFMS at a variety of time intervals. We initially chose trypsin to study early proteolysis of surface-exposed epitopes at neutral pH because of the relatively large number of arginine and lysine residues in most reovirus capsid proteins. For these studies it also was important to maintain viral structural integrity. Preliminary MS experiments tested various buffer solutions. Reovirus virion particles were unstable in ammonium bicarbonate solution (normally suitable for MALDI analysis; data not shown), but virion structure was retained in Tris-HCl buffer with physiologic concentrations of salts (normally too high for MS analyses). To simplify MS analyses, controlled experiments examined the amount of signal noise normally caused by salts in the mass spectra of virus digests by comparing spectra generated from virus alone, protease alone, and virus plus protease in buffers of various ionic strength. In addition, true peptide fragments appeared with increasing intensities during

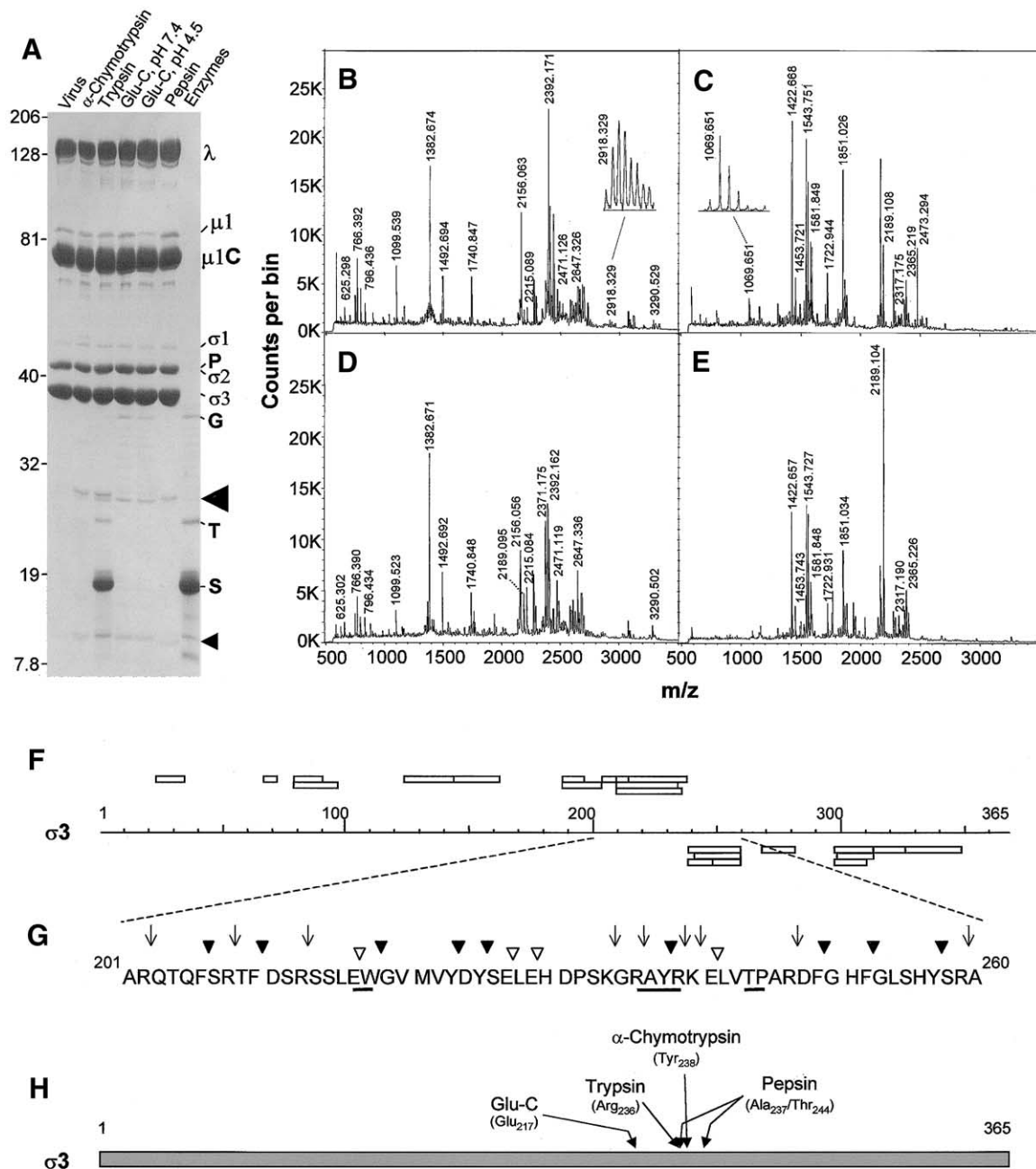


Fig. 2. Identification of large and small σ 3 peptide fragments. (A) Electrophoretic separation of 2.5×10^{11} virions digested with 10 μ g/ml of indicated proteases for 30 s (α -chymotrypsin, trypsin, and pepsin) or for 1.5 h (Glu-C) and resolved in a 6–16% linear gradient SDS-PAGE ($16 \times 16 \times 0.1$ cm) at 6.0 W for 5 h. Molecular weights, determined from coelectrophoresis of Kaleidoscope Rainbow Marker, are indicated to left of gel. Virion proteins, enzymes (~250 ng), and inhibitors (P: pepsin; G: Glu-C; T: trypsin/chymotrypsin; S: soybean trypsin inhibitor), and potential σ 3 digestion products (indicated by arrowheads and subsequently analyzed by mass spectrometry) are indicated to right of gel. (B–E) MALDI-Qq-TOF mass spectra obtained from tryptic in-gel digestions of (B) α -chymotrypsin-generated large fragment (indicated by large arrow head in A), (C) α -chymotrypsin-generated small fragment (indicated by small arrow head in A), (D) trypsin-generated large fragment, and (E) trypsin-generated small fragment. Values above peaks indicate measured m/z values of peaks identified as corresponding to virion protein σ 3 (shown in Table 1). Peak in B with m/z value of 2918.329 (subsequently shown to correspond to amino acid sequence 214–238) and peak in C with m/z value of 1069.651 (subsequently shown to correspond to amino acid sequence 239–247) are expanded in insets. (F) Diagrammatic representation of strategy to determine where each protease initially cleaves virion-associated σ 3. Rectangles above sequence scale correspond to MS-predicted tryptic fragments found in large (~27-kDa polypeptide) and rectangles below sequence scale correspond to MS-predicted tryptic fragments found in small (~12.5-kDa polypeptide) as listed in Table 1. (G) Detailed sequence of σ 3 protein between amino acid residues 201 and 260, indicating potential cleavage sites for trypsin (∇), α -chymotrypsin (\blacktriangledown), and Glu-C (\downarrow). Underline corresponds to cleavage sites identified as initially used. (H) Diagrammatic representation of entire σ 3 protein with initial cleavage sites of tested proteases indicated.

Table 1
Tryptic peptides identified in 27- and 12.5-kDa fragments after brief digestion of $\sigma 3$ with indicated proteases

$\sigma 3$ Sequence assignment	Observed mass					Calculated mass	$\Delta(\text{Da})^a$
	Chymotrypsin	Trypsin	Glu-C, pH 7.4	Glu-C, pH 4.5	Pepsin		
Tryptic fragments found in large peptides							
Tryptic fragments found in all protease digests							
203–208	766.392	766.390	766.386	766.385	766.399	766.385	± 0.014
67–72	796.437	796.434	796.420	796.432	796.437	796.433	± 0.013
187–196	1099.540	1099.537	1099.527	1099.524	1099.548	1099.528	± 0.020
23–34	1382.675	1382.671	1382.651	1382.666	1382.683	1382.659	± 0.024
79–90	1492.701	1492.692	1492.702	1492.687	1492.701	1492.682	± 0.020
187–202	1740.849	1740.848	1740.843	1740.838	1740.850	1740.841	± 0.009
Tryptic fragments found in some protease digests							
209–213	625.298	625.302		625.298	625.303	625.294	± 0.009
237–244					979.571	979.560	± 0.011
144–162	2156.064	2156.059				2156.050	± 0.014
241–259		2189.095 ^b				2189.088	± 0.008
79–96	2215.023	2215.084	2215.075			2215.068	± 0.045
79–97		2371.176	2371.179	2371.165		2371.169	± 0.010
123–143	2392.177	2392.165		2392.164	2392.186	2392.160	± 0.026
214–234	2471.126	2471.119		2471.123	2471.150	2471.107	± 0.043
214–236	2647.326	2647.337		2647.335 ^b	2647.344	2647.329	± 0.015
214–237					2755.289	2755.270	± 0.019
214–238	2918.359					2918.330	± 0.029
209–236	3290.539	3290.502		3290.522 ^b		3290.505	± 0.034
Tryptic fragments found in small peptides							
Tryptic fragments found in all protease digests							
248–259	1422.669	1422.657	1422.668	1422.653	1422.653	1422.655	± 0.014
268–281	1453.751	1453.743	1453.753	1453.746	1453.742	1453.743	± 0.010
313–326	1542.729	1542.727	1542.734	1542.729	1542.714	1542.716	± 0.018
297–310	1581.852	1581.850	1581.820	1581.836	1581.833	1581.838	± 0.018
297–312	1851.037	1851.034	1850.996	1851.019	1851.022	1851.023	± 0.027
327–348	2365.230	2365.226	2365.201	2365.224	2365.215	2365.214	± 0.016
Tryptic fragments found in some protease digests							
239–247	1069.657					1069.648	± 0.009
249–259			1307.633	1307.637	1307.636	1307.628	± 0.009
298–312	1722.944	1722.931	1722.925	1722.935		1722.928	± 0.016
245–259					1746.852	1746.850	± 0.002
241–259	2189.114	2189.107			2189.091	2189.088	± 0.026
240–259	2317.190	2317.190				2317.183	± 0.007
239–259	2473.294					2473.284	± 0.010
238–259					2636.351	2636.348	± 0.003

^a Values represent total range between five treatments.

^b Indicated peak present in very low quantity.

longer digestion times, and individual peaks were subsequently identified by MS/MS measurements. Initial analyses of a complete tryptic digest of reovirus serotype 1 Lang (T1L) virions allowed subsequent identification of the major outer capsid proteins $\sigma 3$ and $\mu 1$, and the core proteins $\lambda 1$, $\lambda 2$, and $\sigma 2$ (data not shown). Identities of all peaks resolved in this initial tryptic digest spectrum were corroborated by tandem MS and used as reference for amino acid assignments in this study.

Representative spectra, which resulted from digestion of 5 mg/ml T1L virus with 5 $\mu\text{g/ml}$ trypsin at pH 7.4 for short

periods of time, are shown in Fig. 3. NCBI and SwissPro database searches of peak masses and subsequent MS/MS analyses of selected peaks against all available viral and nonviral proteins led to the identification of indicated $\sigma 3$ and $\mu 1$ peptides. Results are summarized in Table 2. Examination of peptide fragments released during shorter digestion periods predicted that the first peptide released from $\sigma 3$, at 90 s, with an m/z value of 2684.226, corresponded to residues Ser₂₁₄-Arg₂₃₆ (predicted m/z value of 2684.230) (Table 2A). The low-energy MS/MS spectrum of this ion revealed a set of b and y ions shown in Fig. 4A. The b and

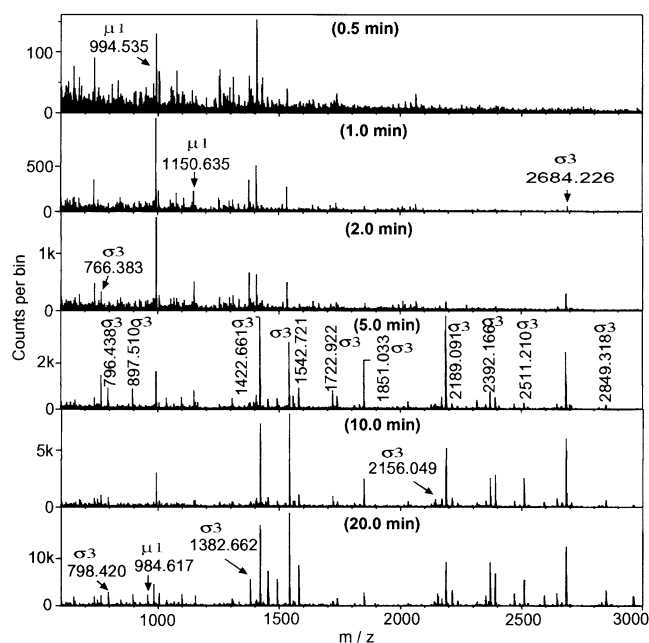


Fig. 3. Peptide mapping of tryptic T1L virion digests. 5 mg/ml purified T1L virions were digested with 5 μ g/ml trypsin at 37°C. At various time intervals, aliquots were collected and analyzed by MALDI-Qq-TOFMS. Each peak was then selected and subjected to MS/MS analysis (see Fig. 4) to identify each peptide fragment (compiled in Table 2A). Each peak is labeled in the time frame at which it is first detected.

y ion “ladder” corresponds to the predicted sequence of the Ser₂₁₄-Arg₂₃₆ fragment. With increased digestion times, additional peptides that corresponded to regions immediately N-terminal of, and C-terminal of the Ser₂₁₄-Arg₂₃₆ peptide, were identified (Fig. 5A and Table 2A) and confirmed by MS/MS (data not shown). Peptides corresponding to virtually the entire σ 3 protein sequence were identified within 10 min of digestion (Fig. 5A and Table 2A), further attesting to the rapid removal of this protein from virions during proteolytic digestion. A peptide corresponding to the N-terminus of protein σ 3 was not initially identified by a direct database search with MS-Tag, but a major unidentified peak at m/z 2511.2 was observed.

The predicted N-terminal tryptic peptide of σ 3 consisting of amino acid residues 1–22 has a calculated m/z value of 2469.202 that was absent in the mass spectrum (Fig. 3). This could indicate that the N-terminal portion of σ 3 remains attached to particles or that this peptide may be modified in such a way that its mass is different from that predicted. The low-energy MS/MS spectrum of the ion at 2511.21 revealed a set of b and y ions shown in Table 3A, corresponding to the N-terminal peptide 1–22 of σ 3. The masses of all b'_n fragment ions and the parent ion (y'_{22}) are 42.010 Da more than the calculated values, indicating the N-terminal methionine residue of σ 3 was modified by acetylation.

Inspection of additional trypsinization time points indicated that the C-terminal peptide of the μ 1/ μ 1C protein at m/z 994.535, corresponding to amino acid residues 695–703 (and confirmed by MS/MS), was released within 30 s of

digestion, 60 s before the detection of the first σ 3 peptide (Figs. 3 and 5A). Other μ 1/ μ 1C peptides were not detected until 20 min after trypsinization, 10 min after σ 3 had been nearly completely digested and removed from particles (Figs. 3, 5A, and Table 2).

Similar experiments also were performed on purified T1L virions with another protease at neutral pH, and the released peptide fragments identified by MALDI-Qq-TOFMS (Table 2 and Fig. 5). Treatment of virions at pH 7.4 with Glu-C, which preferentially cleaves after glutamate residues, resulted in initial liberation of a σ 3 peptide corresponding to Gly₁₉₉-Asp₂₁₇, followed by subsequent release of peptides N-terminal and C-terminal to that peptide. Similar to results obtained with trypsin, treatment of purified reovirus virions with Glu-C resulted in digestion of μ 1 before digestion of σ 3. However, Glu-C digestion initially generated a peptide fragment at m/z 1374.716, which corresponds to residues 513–525 within μ 1 (Table 2 and Fig. 5B). This, as well as all indicated Glu-C peptide fragment assignments (Table 2), were confirmed by MS/MS.

Digestion of σ 3 under acidic conditions

As indicated earlier, there were subtle differences in the identities of some detected tryptic peptides after Glu-C digestion was performed at either pH 7.4 or 4.5 (Table 1, compare columns 4 and 5). Entry of reovirus is thought to involve conformational changes associated with acidic treatment prior to proteolysis. Therefore, we also examined proteolysis of purified virions at low pH. Samples of purified T1L virions were adjusted to pH 3.0 or 4.5 and then digested with pepsin or Glu-C, respectively, at 37°C. Equivalent aliquots were removed at various time points ranging from 30 s to 24 h. Digestions were inhibited with the immediate addition of 10 mM Tris pH 7.4 to pepsin-digested samples (confirmed to raise the pH sufficiently to irreversibly inhibit the enzyme), or by the addition of 0.1% trifluoroacetic acid (TFA) to Glu-C-digested samples. Digestion products were analyzed by MALDI-Qq-TOFMS. All Glu-C peptides were identified by peptide mapping and high-accuracy mass measurements to allow direct comparison to corresponding peaks previously identified by peptide mapping and MS/MS measurements of the pH7.4 Glu-C-treated samples (above), whereas all identified pepsin fragments were sequenced with MS/MS analysis because of pepsin's nonspecific cleavage. For example, the low-energy MS/MS spectrum of the second ion detected after pepsin digestion (at m/z 1419.741) revealed a set of b and y ions shown in Table 3B, corresponding to amino acids 195–206. Results of both acidic digestions are summarized in Table 2C. The first peak identified after digestion of intact virions with Glu-C at pH 4.5 corresponds to σ 3 amino acids Gly₁₉₉-Asp₂₁₇, the same peptide initially released by this protease at pH 7.4. However, the first peptide identified after pepsin digestion of virions at pH 3.0 corresponded to amino acids Asn₁₁₁-Leu₁₂₈ in σ 3. This pepsin-digested fragment is about

Table 2

Peptide fragments generated from a time-course digestion of T1L virions

Time (min)	$\sigma 3$	m/z (Obs.)	m/z (Calc.)	μl	m/z (Obs.)	m/z (Calc.)
A: Tryptic fragments (pH 7.4) ^a						
0.5	—	—	—	695–703	994.535	994.532
1.0	—	—	—	695–704	1150.635	1150.633
1.5	214–236	2684.226	2684.230	—	—	—
2.0	203–208	766.383	766.385	—	—	—
2.5	—	—	—	—	—	—
3.0	209–213	625.290	625.295	—	—	—
	248–259	1422.661	1422.655	—	—	—
	241–259	2189.091	2189.089	—	—	—
5.0	1–22 ^b	2511.210	2469.202	—	—	—
	67–72	796.438	796.433	—	—	—
	79–90	1492.692	1492.682	—	—	—
	91–97	897.510	897.506	—	—	—
	79–97	2371.167	2371.170	—	—	—
	73–97	3113.515	3113.524	—	—	—
	98–122	2849.318	2849.321	—	—	—
	123–143	2392.166	2392.161	—	—	—
	187–196	1099.538	1099.528	—	—	—
	187–202	1740.846	1740.842	—	—	—
	209–236	3290.496	3290.506	—	—	—
	240–259	2317.191	2317.184	—	—	—
	241–259	2189.091	2189.089	—	—	—
	248–259	1422.661	1422.655	—	—	—
	268–281	1455.686	1455.680	—	—	—
	297–310	1581.841	1581.839	—	—	—
	297–312	1851.033	1851.024	—	—	—
	298–312	1722.922	1722.929	—	—	—
	313–326	1542.721	1542.716	—	—	—
10.0	23–63	4326.029	4326.031	—	—	—
	35–63	2962.389	2962.390	—	—	—
	98–120	2594.166	2594.151	—	—	—
	144–162	2156.049	2156.051	—	—	—
	327–365	4199.064	4199.083	—	—	—
20.0	282–287	798.420	798.422	235–243	984.617	984.621
	23–34	1382.662	1382.659	—	—	—
	214–234	2471.119	2471.108	—	—	—
25.0	163–186	2873.360	2873.368	—	—	—
B: Glu-C digested fragments (pH 7.4) ^a						
0.5	—	—	—	—	—	—
1.0	—	—	—	—	—	—
1.5	—	—	—	—	—	—
2.0	—	—	—	—	—	—
2.5	—	—	—	513–525	1374.728	1374.716
3.0	—	—	—	—	—	—
5.0	199–217	2188.044	2188.038	—	—	—
10.0	—	—	—	—	—	—
20.0	228–241	1685.878	1685.872	—	—	—
	230–241	1443.750	1443.746	—	—	—
30.0	21–30	1109.565	1109.559	137–153	1923.929	1923.916
50.0	3–20	1995.994	1995.996	—	—	—

Table 2 (continued)

Time (min)	Glu-C (pH 4.5)			Pepsin (pH 3.0) ^a		
	$\sigma 3$	m/z (Obs.)	m/z (Calc.)	$\sigma 3$	m/z (Obs.)	m/z (Calc.)
C: Glu-C and pepsin digested fragments (pH 4.5 and 3.0)						
0.5	—	—	—	—	—	—
1.0	—	—	—	—	—	—
1.5	—	—	—	111–128	1958.942	1958.942
2.0	—	—	—	195–206	1419.741	1419.734
2.5	—	—	—	142–154	1573.778	1573.776
				142–159	2115.057	2115.051
3.0	—	—	—	27–42	1763.804	1763.791
	—	—	—	245–265	2343.204	2343.200
	—	—	—	295–309	1708.903	1708.913
	—	—	—	343–357	1735.783	1735.789
5.0	199–217	2188.036	2188.038	Too many		
10.0	—	—	—			
20.0	228–241	1685.873	1685.872			
	230–241	1443.740	1443.746			
30.0	21–30	1109.550	1109.559			
50.0	3–20	1995.994	1995.996			
60.0	—	—	—			
120.0	176–198	2619.241	2619.245			
	301–309	998.464	998.470			

^a Assignments of all peaks corroborated by tandem MS.^b Acetylation at the N-terminus.

100 residues closer to the N-terminus than where trypsin and Glu-C originally cleaved at pH values of 4.5 and 7.4, suggesting a conformational change in $\sigma 3$ had taken place as a result of the more acidic treatment (Table 2 and Fig. 5).

To determine whether the different peptides released by proteolysis at pH 3.0 were a result of virus dissociation at low pH, purified virions were adjusted to pH 3.0, main-

tained at that pH for various periods of time up to 1 h, and examined by electron microscopy. Reovirus virions maintained at pH 3.0 for as much as 60 min appeared fully intact (data not shown), in agreement with previous data showing no change in morphology and no loss of infectivity after exposure of virions to pH 2.0 (Stanley, 1967; Fields and Eagle, 1973). This suggests that release of a different initial peptide after proteolysis at pH 3.0 as compared to proteolysis at higher pH values is not the result of acid-induced global viral capsid disassociation.

Structural localization of $\sigma 3$ peptides in virions

The available X-ray crystallographic structures of the $\sigma 3$ dimer (Olland et al., 2001) and $\sigma 3/\mu 1$ heterohexamer (Liemann et al., 2002) were analyzed and manipulated with the computer-based programs RasMol 2.7.1. (Bernstein and Sons, Belpport, NY), PyMOL (DeLano, 2002), and Molscript (Kraulis, 1991) to determine where each of the tested proteases initially cut $\sigma 3$ within the virus and where each initially released peptide is located. The initial α -chymotrypsin cleavage site (Tyr₂₃₈), trypsin cleavage site (Arg₂₃₆), and pepsin cleavage sites (Ala₂₃₇) are located most distally in the $\sigma 3$ protein (Fig. 6A, C, and D) at the top of the $\sigma 3/\mu 1$ heterohexamer (Fig. 7). In addition, pepsin fragment Asn₁₁₁-Leu₁₂₈ is located at the top of the molecule but constitutes a separate group of amino acids (Figs. 6, 7). Glu-C fragment Gly₁₉₉-Asp₂₁₇ is located on a different surface of the protein and the initial site cut by Glu-C (Asp₂₁₇) is located well down the side of the $\sigma 3$ molecule (Figs. 6 and 7).

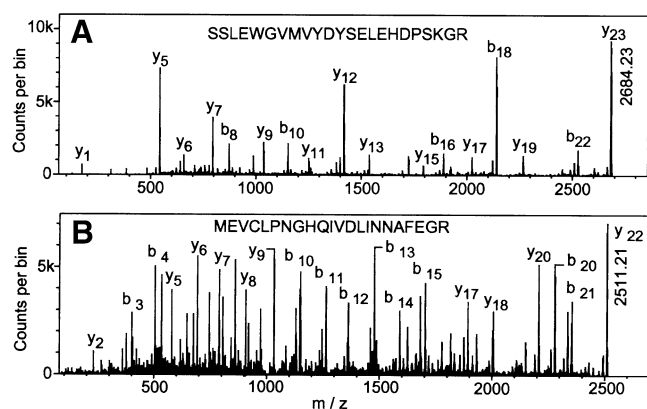


Fig. 4. MS/MS spectra of selected tryptic ions. (A) The mass spectrum of the first tryptic ion detected, at m/z 2684.23, was acquired at a laser repetition rate of 7 Hz. The ion was dissociated using argon as the collision gas. The collision energy was set to 140 eV and then slightly adjusted to obtain optimum fragmentation of the parent ion. Spectra data acquisition was performed using software developed in-house (Tofma, University of Manitoba, MB). Sequence analysis indicates the ion corresponds to $\sigma 3$ peptide 214–236. (B) The mass spectrum of the ion at m/z 2511.21 was acquired and dissociated as above. Sequence analysis indicates the N-terminal acetylation of $\sigma 3$ peptide 1–22.

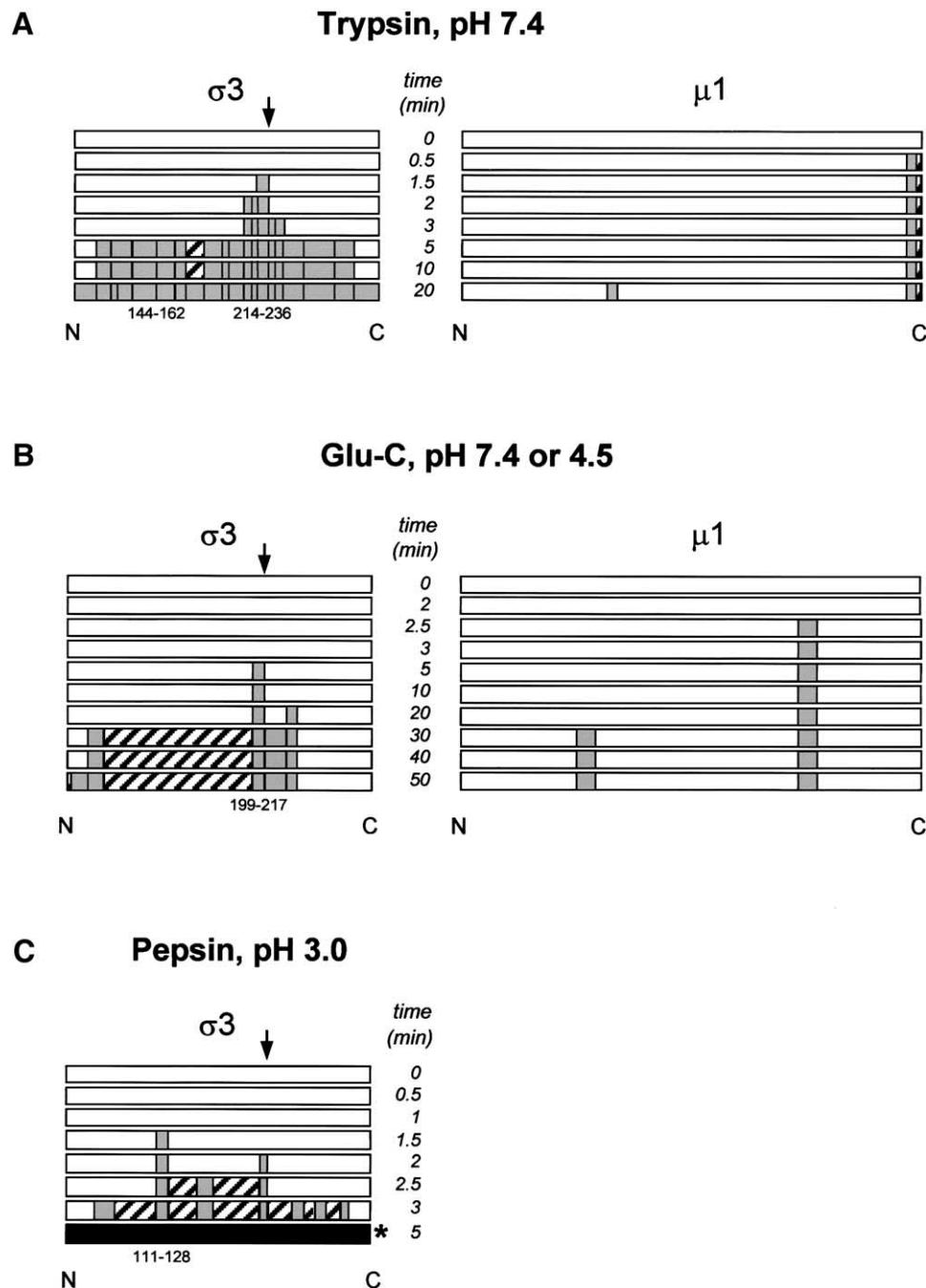


Fig. 5. Diagrammatic representation of time-course enzymatic digests of $\sigma 3$ and $\mu 1$ outer capsid proteins. (A) Time-course digestion of T1L $\sigma 3$ and $\mu 1$ by trypsin at pH 7.4. (B) Time-course digestion of T1L $\sigma 3$ and $\mu 1$ by Glu-C at pH 7.4 or 4.5 (results identical at both pH values). (C) Time-course digestion of T1L $\sigma 3$ by pepsin at pH 3.0. White bar: undigested protein; gray bar: peptides identified by MS or MS/MS; hatched bar: nonidentified peptides; black bar: too many fragments to identify. Initial peptides cleaved are indicated with numbers below bars, with the exception of 144–162 in A (see Discussion).

Discussion

Characterization of virus proteins by MALDI-Qq-TOFMS

The recent development of orthogonal-injection MALDI-TOF provides a powerful tool for the structural characterization of virus proteins. Compared to electrospray ionization mass spectrometry, the tolerance to impurities permits the sample to be analyzed without any further sep-

aration or purification by methods similar to liquid chromatography. This alleviates sample losses associated with additional separation, which was particularly important for observation of the first peptides released during time-course digestion studies, which appeared after a few seconds at relatively low concentration compared to the salts. The technique is also fast, even with a low repetition-rate laser, allowing a large number of samples to be acquired in time-course studies. This is essential to distinguish changes in the

Table 3
Comparison of MS/MS fragmentation of selected ions with calculated mass values

b ion	<i>m/z</i> (Obs.)	MH ⁺ (Calc.)	Δ (Da)	y ion	<i>m/z</i> (Obs.)	MH ⁺ (Calc.)	Δ (Da)
A: MS/MS comparison of tryptic ion at <i>m/z</i> 2511.21 with mass values calculated assuming an acetylated N-terminus indicates residues 1–22: MEVCLPNGHQIVDLINNAFEGR							
b'1	—	174.059	—	y1	175.116	175.120	−0.004
b'2	303.098	303.101	−0.003	y2	232.140	232.141	−0.001
b'3	402.170	402.170	0.000	y3	361.180	361.184	−0.004
b'4	505.179	505.179	0.000	y4	—	508.252	—
b'5	618.272	618.263	+0.009	y5	579.293	579.289	+0.004
b'6	—	715.316	—	y6	693.332	693.332	0.000
b'7	—	829.359	—	y7	807.371	807.375	−0.004
b'8	886.387	886.380	+0.007	y8	920.458	920.459	−0.001
b'9	1023.439	1023.439	0.000	y9	1033.541	1033.543	−0.002
b'10	1151.488	1151.498	−0.010	y10	1148.565	1148.570	−0.005
b'11	1264.581	1264.582	−0.001	y11	1247.636	1247.638	−0.002
b'12	1363.661	1363.650	+0.011	y12	1360.724	1360.723	+0.001
b'13	1478.672	1478.677	−0.005	y13	1488.775	1488.781	−0.006
b'14	1591.769	1591.761	+0.008	y14	1625.828	1625.840	−0.012
b'15	1704.854	1704.845	+0.009	y15	1682.871	1682.862	+0.009
b'16	1818.902	1818.888	+0.014	y16	—	1796.905	—
b'17	1932.936	1932.931	+0.005	y17	1893.946	1893.957	−0.011
b'18	2003.974	2003.969	+0.005	y18	2007.027	2007.041	−0.014
b'19	2151.048	2151.037	+0.011	y19	2110.040	2110.051	−0.011
b'20	2280.074	2280.079	−0.005	y20	2209.119	2209.119	0.000
b'21	2337.108	2337.101	+0.007	y21	—	2338.161	—
b'22	2493.205	2493.202	+0.003	y'22	2511.210	2511.212	−0.002
B: MS/MS comparison of pepsin ion at <i>m/z</i> 1419.74 indicates residues 195–206: VRLEGDARQTQF							
b1	—	100.076	—	y1	—	166.087	—
b2	256.178	256.177	0.001	y2	—	294.145	—
b3	369.257	369.261	−0.004	y3	—	395.193	—
b4	498.301	498.304	−0.003	y4	—	523.252	—
b5	—	555.326	—	y5	679.354	679.353	0.001
b6	670.351	670.352	−0.001	y6	750.393	750.390	0.003
b7	—	741.390	—	y7	—	865.417	—
b8	897.495	897.491	0.004	y8	922.425	922.438	−0.013
b9	1025.556	1025.549	0.007	y9	—	1051.481	—
b10	1126.591	1126.597	−0.006	y10	1164.561	1164.565	−0.004
b11	1254.649	1254.656	−0.007	y11	—	1320.666	—
b12	1401.716	1401.724	−0.008	y12	1419.738	1419.734	0.004

spectrum to discriminate against noise. The high mass accuracy (average mass errors are 0.01 Da or less (Tables 1–3)) in both single MS and MS/MS mode gives high success rate in protein identification by database searches. The high accuracy in MS/MS measurements is also particularly useful for de novo peptide sequencing or for identification of posttranslational modifications.

Previous work has indicated that virtually all reovirus proteins are blocked at their N-termini (Pett et al., 1973). Outer capsid protein $\mu 1$ contains a myristoyl group at its N-terminus (Nibert et al., 1991). Very recent reports also have identified an acetyl group at the N-terminus of $\sigma 3$ (Jane-Valbuena et al., 2002), as confirmed in this study (Table 3A), and at the amino-terminus of minor reovirus core protein $\mu 2$ (Swanson et al., 2002). The function of an acetyl group at the N-terminus of $\sigma 3$ is unknown, but, based on roles of other acetylated molecules, may serve to provide

structural stability (Cumberlidge and Isono, 1979; Nguyen et al., 2000) and/or promote protein complex assembly (Berger et al., 1981; Tercero and Wickner, 1992). Another possibility is that the acetylation of its N-terminus may be required to differentiate the role(s) $\sigma 3$ plays during the virus life cycle. Protein $\sigma 3$ binds dsRNA (Huisman and Joklik, 1976), which may occur only when the protein dimerizes (Olland et al., 2001), $\sigma 3$ is thought to counteract the reovirus induced host antiviral defence mechanism (Yue and Shatkin, 1997), and $\sigma 3$ may play a role in the assortment complex during particle assembly (Antczak and Joklik, 1992). Previous studies also have shown that the efficiency of cellular translation is determined by the cytoplasmic level of $\sigma 3$ that is not complexed with $\mu 1$ (Schmechel et al., 1997). Thus, some level of regulation must dictate which of the numerous roles $\sigma 3$ functions in. One possible means of control is through $\mu 1$ binding, which would effectively

remove freely available $\sigma 3$ protein. Another regulatory role has been postulated to occur upon $\sigma 3$ dimerization (Olland et al., 2001). A third possibility, currently under investigation, is that acetylation of $\sigma 3$ may influence its regulatory roles, as suggested for a number of other acetylated proteins (reviewed in Polevoda and Sherman, 2000).

Morphology and orientation of reovirus outer capsid proteins

To date, there has been only limited evidence as to the exact placement of viral proteins within the bilayered capsids of reovirus. The high-resolution crystal structure of the innermost capsid core is known (Reinisch et al., 2000). In addition, the high-resolution crystal structure of outer capsid protein $\sigma 3$ dimers (Olland et al., 2001) and $\mu 1/\sigma 3$ heterohexamers (Liemann et al., 2002) have been determined and fitted, along with the core crystal structure, into cryo-EM models of the entire virion (Liemann et al., 2002). A large number of dramatic conformational changes, thought to be important in different stages of reovirus disassembly, have been described. These include changes during cell attachment (Fernandes et al., 1994), uncoating (Dryden et al., 1993), and transcription (Yeager et al., 1996). However, the entire virion has, to date, resisted crystallization and environmentally induced conformational alterations in the complex, that may result from exposure to acidic pH and pepsin (such as when this enteric virus passes through the stomach where the pH can be as low as 0.8; Vander et al., 1994), are poorly understood. The present analyses of purified virions by limited proteolysis and mass spectrometry has furthered this understanding and has provided evidence about the nature of viral outer capsid proteins in terms of structure, orientation, and conformational alterations.

Digestion of reovirus virions generally resulted in sequential digestion of outer capsid protein $\sigma 3$, followed by sequential digestion of $\mu 1$. When virions were digested with α -chymotrypsin, pepsin, trypsin, and Glu-C protease, $\sigma 3$ digestion initiated within a protease hypersensitive region (Fig. 2), which indicates this region is located on the external surface of the virion. Other studies that examine $\sigma 3$ digestion have recently determined that this hypersensitive region is also initially cut by other proteases, including alkaline proteases such as trypsin (Jane-Valbuena et al., 2002) and acidic proteases, such as cathepsins (Ebert et al., 2002; Jane-Valbuena et al., 2002). Our strategy of directly measuring peptides released from protease-treated intact particles by MALDI-Qq-TOFMS also allowed us to determine subsequent proteolytic events during the cascade of $\sigma 3$ and $\mu 1$ digestion. These time-course studies showed that protein $\sigma 3$ digestion proceeded bidirectionally toward both the N-terminus and the C-terminus, except that tryptic peptide fragment Ser₁₄₄-Lys₁₆₂ did not appear until after the remainder of the protein had been completely digested, which suggests this peptide may have been particle bound,

and/or it may be this region of $\sigma 3$ that interacts with underlying $\mu 1$, as suggested in the $\mu 1/\sigma 3$ heterohexamer structure (Liemann et al., 2002). This result was not observed when virions were digested with Glu-C protease, which may be attributable to the fact that there are far fewer Glu-C cleavage sites that could result in the lack of detection of such a fragment.

Despite the belief that $\sigma 3$ digestion appears to precede $\mu 1$ digestion (Fig. 1; see also, for example, Nibert et al., 1995), digestion of intact virions with trypsin initially released a short fragment of eight amino acid residues which corresponds to the extreme C-terminus of $\mu 1$, before any $\sigma 3$ peptides were released. The appearance of this $\mu 1$ fragment suggests that the C-terminus of $\mu 1$ is accessible to trypsin and hence may be exposed on the surface of the virion, consistent with a recent hypothesis that this unresolved portion of $\mu 1$ in the $\mu 1/\sigma 3$ crystal structure constitutes an exposed blob within the virus (Liemann et al., 2002). Furthermore, immunologic assays, using monoclonal antibodies directed against $\mu 1$, react with intact virions, which suggest a portion of $\mu 1$ is exposed on the virion surface (Hayes et al., 1981; Virgin et al., 1994), as also suggested by other structural studies (Dryden et al., 1993; Liemann et al., 2002). Initial digestion of $\mu 1$ by such proteases may be a prerequisite for reovirus uncoating and membrane penetration, a possibility that is currently under study.

Possible pH-induced conformation changes within the outer viral capsid

Viruses, such as reovirus, maintain high structural stability during the extracellular portions of their replicative cycles, yet are significantly dynamic during other stages in their life cycle, namely during cell attachment (Fernandes et al., 1994), uncoating (Dryden et al., 1993), and transcription (Yeager et al., 1996). Mass spectrometry has been used in the past to provide evidence of conformational changes in other viruses such as in flock house virus (Bothner et al., 1998), human rhinovirus (Lewis et al., 1998), and in Sindbis virus (Phinney et al., 2000). Flock house virus capsids underwent a change in conformation when treated with different solvents (Bothner et al., 1998). Binding of antiviral agents to human rhinovirus caused localized conformational changes that stabilized the viral capsid (Lewis et al., 1998). Low pH exposure caused conformational changes in the membrane glycoproteins of Sindbis virus (Phinney et al., 2000).

The present MALDI-Qq-TOFMS analyses of purified reovirus virions identified different initially released peptides when T1L was digested with different proteases under different pH conditions. This may provide evidence that T1L $\sigma 3$ molecules undergo a conformational change when exposed to acidic pH. However, we acknowledge difficulty in interpreting data from using two different enzymes with different amino acid cleavage specificities under these significantly different pH condi-

tions. Unfortunately, we are not aware of any single enzyme that operates efficiently under both pH conditions. Thus, it also is possible the different initially released peptides result from different enzyme specificities. To attempt to alleviate the difficulty in interpretation of experiments carried out with different enzymes under different pH conditions, we also performed experiments with Glu-C, which, unlike trypsin and pepsin, can operate under widely different pH conditions (pH 7.4 and 4.5). Tryptic digests of $\sigma 3$ proteins that had been cut only once with Glu-C (Fig. 2 and Table 1, columns 4 and 5) revealed minor differences, while kinetic studies of the cascade of proteolytic events in whole particles (Fig. 5, and compare Table 2B to 2C) revealed no differences. Part of the difficulty with using Glu-C for such comparative studies is the lower frequency of appropriate cleavage sites in $\sigma 3$ and their uneven distribution, which results in some sequence-predicted peptides that would lie outside the normal operating m/z range (500–5000) of the MALDI-Qq-TOFMS used. Other apparent paradoxes are that the initial Glu-C cleavage site (Glu₂₁₇) and initial fragment released by Glu-C digestion (Gly₁₉₉-Glu₂₁₇) appear, predicted from fitting the $\sigma 3/\mu 1$ heterohexamer crystal structure into whole particle cryo-EM reconstructions (Liemann et al., 2002), to reside in regions of the $\sigma 3$ protein that abut other $\sigma 3$ proteins located in adjacent $\sigma 3/\mu 1$ heterohexamers. Resolution of this apparent conflict, which may result from the different methodologies of comparing static crystal structures to virions digested in solution, the latter of which may be capable of “breathing,” as well as additional studies to better examine possible pH-induced conformational alterations, are underway. Interestingly, despite these differences, $\sigma 3$ is completely cleaved rapidly under each of these conditions to generate a subviral particle, which suggests the virus has evolved to successfully uncoat itself under a variety of conditions likely to be encountered during infection.

In conclusion, our results pinpoint those regions of the reovirus outer capsid protein $\sigma 3$ that are initially cut by a variety of proteases, and thus presumably exposed in virions at a variety of pH values, confirm the presence of an acetyl group at the amino-terminus of this protein in virions, and delineate the cascade of $\sigma 3$ cleavages as the virion is converted into an ISVP.

Materials and methods

Cells and viruses

T1L was plaque purified and grown in mouse L929 cell monolayers in Joklik-modified minimal essential medium (GIBCO, Grand Island, NY) supplemented to contain 2.5% fetal calf serum (Gibco-BRL Life Technologies, Burlington, Ontario), 2.5% VSP neonate bovine serum (Biocell, Carson,

CA), 2 mM glutamine, 100 U penicillin per milliliter, 100 μ g streptomycin sulfate per milliliter, and 1 μ g amphotericin-B per milliliter, as previously described (Coombs et al., 1994). Large amounts of virus were grown in spinner culture, extracted with Vertrel-XF, and purified in cesium chloride gradients as described (Mendez et al., 2000). Virus bands were harvested, dialyzed extensively against dialysis buffer (D buffer: 150 mM NaCl, 15 mM MgCl₂, 10 mM Tris, pH 7.4), and stored at 4°C until used. Protease digestion analyses were conducted multiple times on virions that had been prepared and purified on three separate occasions.

Proteolysis

Endoproteases Glu-C (*Staphylococcus aureus* V8) and pepsin (Boehringer Mannheim GmbH) were resuspended as stock solutions (0.1 mg/ml) in appropriate buffers and stored at –20°C. Chymotrypsin and sequencing-grade trypsin (Sigma Chemical Co.) solutions were freshly prepared in D buffer (pH 7.4). Enzyme activities were verified with standard protein BSA (Sigma). Aliquots of virus were digested with proteases within 14 days of having been dialyzed. Purified virions were usually diluted to final concentrations of 3–5 mg/ml into buffers precalibrated to produce final pH values of 3.0, 4.5, or 7.4, and then the diluted virions were incubated at 37°C with various concentrations of Glu-C, chymotrypsin, trypsin, or pepsin at a final protein-to-enzyme ratio of 10:1 to 1000:1. Under conditions of high viral particle concentration, extended proteolysis of virions was confirmed to generate core particles (data not shown). Aliquots from the digestion reactions were collected at various time intervals. For peptides subsequently resolved in polyacrylamide gels, further digestion of viral proteins was inhibited by addition of phenylmethylsulfonyl fluoride to chymotrypsin digestions to a final concentration of 5 mM, by addition of soybean trypsin inhibitor to a final concentration of 125 μ g/ml to trypsin digestions, by addition of 1/25th volume of 1 M Tris pH 8.8 to pepsin reactions, and by chilling Glu-C reactions. One-quarter volume of 5× electrophoresis sample buffer was added to each reaction and immediately boiled for gel analysis (below). Further digestion of peptides to be directly measured by MS was inhibited by addition of an equal volume of 0.1% trifluoroacetic acid to samples digested with chymotrypsin, trypsin, or Glu-C, or by addition of an equal volume of 10 mM Tris pH 7.4 to samples digested with pepsin. For direct MS analyses, nascent digests were immediately placed on ice to arrest proteolysis and diluted to 1 mg/ml with dH₂O. A 1:1 ratio of diluted sample and 2,5-dihydroxybenzoic acid (DHB) matrix solution was prepared and 0.5 μ l was placed onto a MALDI sample plate and dried for MALDI-Qq-TOFMS analysis. Capacity of the respective treatments to arrest specific proteases was confirmed in time-course digestions or with standard BSA.

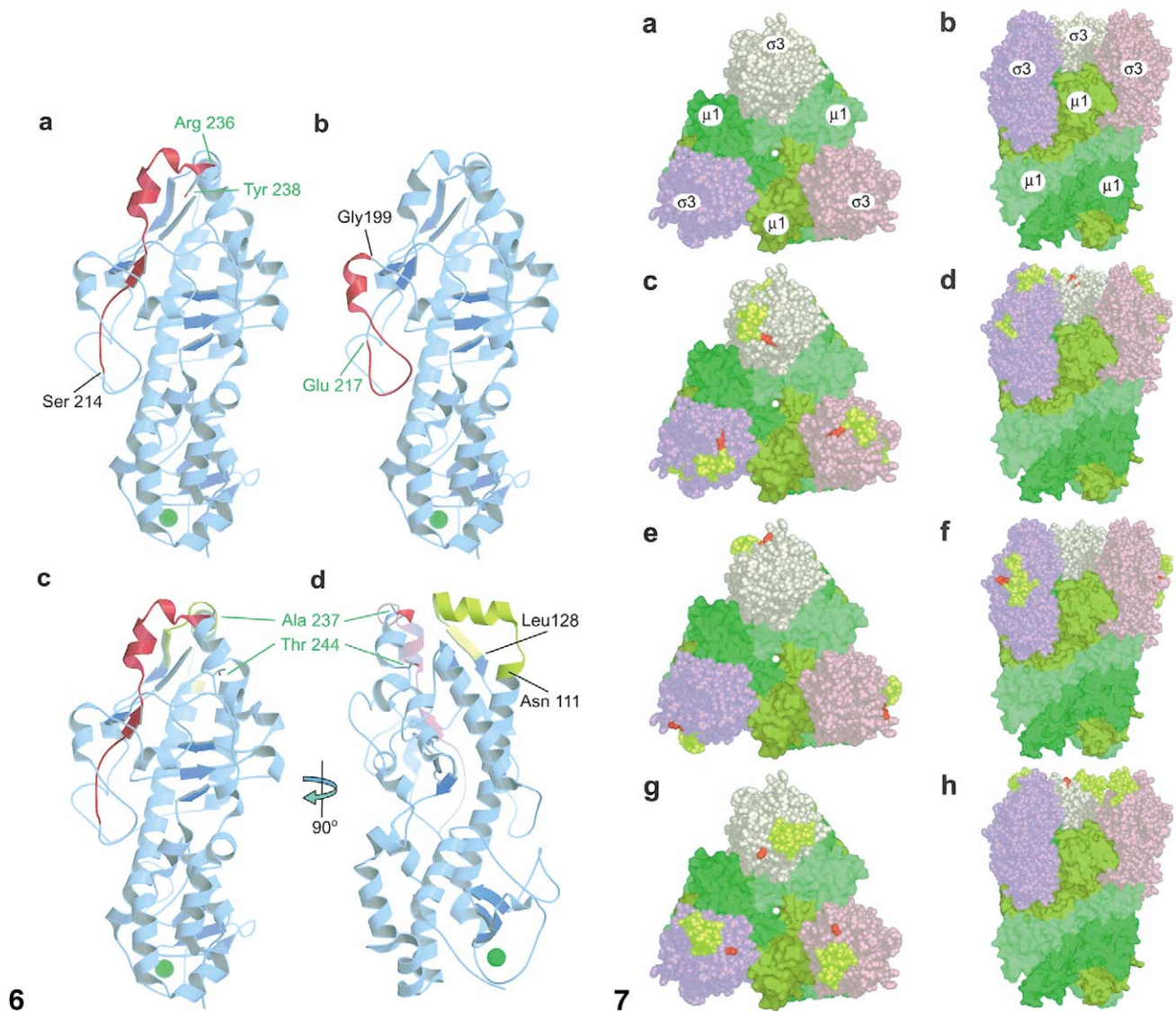


Fig. 6. X-ray crystallographic structure of the $\sigma 3$ monomer. The orientation of the $\sigma 3$ monomer (adapted and modified from Olland et al., 2001) was examined with Molscript (Kraulis, 1991) and rendered with Raster3D (Merritt and Bacon, 1997). With the exception of selected regions shown in red (A–D) or olive (C–D), helices and turns are in light blue and β -sheets are in dark blue. (A) The selected red region corresponds to residues Ser₂₁₄-Arg₂₃₆, the initial tryptic fragment at pH 7.4. (B) The selected red region corresponds to residues Gly₁₉₉-Glu₂₁₇, the initial Glu-C protease fragment at pH 7.4. (C) and (D) The red region corresponds to residues Ser₂₁₄-Arg₂₃₆ from trypsin digestion at pH 7.4 and the olive region corresponds to residues Asn₁₁₁-Leu₁₂₈ from pepsin digestion at pH 3.0. In (D) the $\sigma 3$ monomer is rotated 90° clockwise about the y-axis. The large green dot near the bottom of each image corresponds to the previously determined zinc ion (Olland et al., 2001).

Fig. 7. X-ray crystallographic structure of the $\sigma 3/\mu 1$ heterohexamer. The orientation of the $\sigma 3/\mu 1$ heterohexamer (adapted and modified from Liemann et al., 2002) using PyMol (DeLano, 2002) is shown in space-filling mode. Panels (A), (C), (E), and (G) show top views and panels (B), (D), (F), and (H) show side views of the structure. Individual proteins are identified in (A) and (B); $\mu 1$ are depicted in forest, olive, and lime; $\sigma 3$ are depicted in wheat, pink, and salmon. Initial cleavage sites (red residues) and entire peptide fragments initially released from $\sigma 3$ during time-course digestions (denoted in yellow) are indicated for trypsin (C, D), Glu-C (E, F), and pepsin (G, H).

Sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE)

Proteolytically digested samples were resuspended in electrophoresis sample buffer (240 mM Tris–HCl, pH 6.8, 1.5% dithiothreitol, 1% SDS), heated to 95°C for 5 min, and resolved in various gradient SDS–PAGE gels, using the discontinuous Tris/glycine buffer system (Laemmli, 1970).

Gels were fixed and stained with Coomassie brilliant blue R-250.

In-gel digestion and extraction of peptides

The following steps were modified from previously published procedures (Rosenfeld et al., 1992; Hellman et al., 1995; Shevchenko et al., 1996; Wilm et al., 1996). The

excised gel bands (from SDS–PAGE gels above) were fragmented and destained by repeated washings with alternating 100 mM NH_4HCO_3 and a 1:1 (v/v) mixture of CH_3CN :100 mM NH_4HCO_3 solutions. After washes, gel pieces were dried in a SpeedVac and reswollen with 20 μl of 25 mM NH_4HCO_3 that contained 30 ng trypsin (Sigma, sequencing grade) for 45 min at 4°C. An additional 200 μl of the buffer was added to cover the gel pieces, which were then incubated at 37°C overnight. The digestion mixture was deactivated with 10 μl 2% TFA and the resultant peptide fragments were eluted by 30 min of sonication in the presence of three successive changes in 150 μl 0.1% TFA, 150 μl 1:1 mixture of acetonitrile:0.1% TFA (v/v), and 150 μl 100% acetonitrile. The whole extraction procedure was repeated three additional times and the collected solutions that contained extracted peptides were pooled and dried.

Mass spectrometry and data analyses

Mass spectra of proteolytic peptides, and tandem mass spectra (MS/MS) of their daughter ions, were obtained on a prototype tandem quadrupole/TOF mass spectrometer (QqTOF) (Shevchenko et al., 1997) coupled to a MALDI source (Chernushevich et al., 1999). Samples were irradiated with a nitrogen laser (Laser Science Inc., model 337ND) operating at a repetition rate of 7 Hz for single MS and 15 Hz for tandem MS. Single MS spectra were acquired in approximately 1 min; tandem mass spectra were accumulated for several minutes. For tandem mass spectrometry, selected ions were fragmented in a collision cell that contained argon. The collision energy was adjusted manually between 30 and 170 eV to obtain optimum fragmentation of the parent ions.

Mass spectra were recorded and analyzed using software (*Tofma*) developed in-house. Two-point, external calibration was performed using a mixture of dalargin and melittin. The peptides and proteins were identified by searching on-line databases against monoisotopic masses using *MS-Tag* and *MS-Fit* (<http://prospector.ucsf.edu/>; <http://prowl.rockefeller.edu/cgi-bin/ProFound>). For comparison, fragment masses from either enzymatic digestions or collision-induced dissociation (CID) were calculated using *ProMac* (Sciex, Concord, ON).

Protein structure analyses

The dimeric $\sigma 3$ X-ray crystallographic structure (Accession No. PDB ID 1FN9, originally determined by Olland et al., 2001) and $(\sigma 3\mu 1)_3$ heterohexameric X-ray crystallographic structure (Accession No. PDB ID 1JMU, originally determined by Liemann et al., 2002) were manipulated and modified with the graphics visualization computer programs RasMol 2.7.1. (Bernstein and Sons, Belpport, NY), PyMol (DeLano, 2002), and Molscript (Kraulis, 1991).

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